Claims

- 1. A nucleic acid sequence comprising any one of the nucleic acid sequences of SEQ ID NOs 1-20, or a subfragment nucleic acid sequence derived from any one
 5 of the sequences of SEQ ID NOs 1-20, wherein an mRNA molecule comprising said sequence has RNA binding protein (RBP) binding activity or regulates the
 - 2. The nucleic acid sequence of claim 1, wherein said subfragment nucleic acid sequence is optimized.

functionality of said/mRNA.

- 3. The nucleic acid sequence of claim 1, wherein the regulation of mRNA functionality comprises an alteration in pre-mRNA processing or in the stabilization, translational efficiency, localization, sequestration, editing, or splicing functions of said mRNA.
- A method of identifying an optimized subfragment of any one of the parent nucleic acid sequences of SEQ ID NOs 1-20, said method comprising isolating a subfragment nucleic acid sequence from said parent nucleic acid sequence,
- assaying RNA molecules comprising said subfragment for RBP binding activity or mRNA functionality, and identifying a subfragment nucleic sequence that maintains an RBP binding activity and/or mRNA functionality that is equivalent to said parent sequence.
- 25 5. The method of claim 4, wherein said subfragment nucleic acid sequence is isolated by restriction enzyme digestion.

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6. The method of claim 4, wherein said subfragment is identified by deletion mapping.

7. The method of claim 4, wherein said mRNA functionality comprises an alteration in pre-mRNA processing or in the stabilization, translational efficiency, localization, sequestration, editing, or splicing functions of said mRNA.

8. A nucleic acid sequence identified as an optimized subfragment of any one of SEQ ID NOs 1-20 by the method of claim 4.

9. A method of identifying a candidate compound having an effect on an RNA/RBP binding pair interaction or mRNA functionality, said method comprising contacting an RNA molecule comprising at least one nucleic acid sequence of any one of SEQ ID NOs 1-20, or at least one optimized subfragment sequence derived from any one of SEQ ID NOs 1-20, with at least one RBP, and at least one test compound, and measuring said RNA/RBP binding pair interaction and/or mRNA functionality, wherein a candidate compound is identified as a test compound that affects said interaction and/or functionality.

20 10. The method of claim 9, wherein said mRNA functionality comprises an alteration in pre-mRNA processing or in the stabilization, translational efficiency, logalization, sequestration, editing, or splicing functions of said mRNA.

11. A method for identifying an RBP that interacts with an RNA molecule comprising the nucleic acid sequence of any one of SEQ ID NOs 1-20, or an optimized subfragment sequence of any one of SEQ ID NOs 1-20, said method

comprising contacting said RNA molecule with at least one RBP, and measuring RNA/RBP binding pair interactions, wherein detection of said interactions identifies said RBP that interacts with said RNA molecule.

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